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Co/Cu-MFF derived mesoporous ternary metal oxide microcubes for enhancing the catalytic activity of the CO oxidation reaction†

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Metal-organic framework (MOF)-based derivatives with uniform micro/mesoporous structures have attracted a great deal of interest in various research fields. Herein, we report a simple strategy to design functional mesoporous ternary metal oxides with controlled composition through direct pyrolysis of Co/Cu bimetal-formate frameworks (Co/Cu-MFFs), which were prepared by a facile one-step liquid-phase precipitation method, exhibiting uniform distribution of two different metal species and good structural integrity. The obtained mesoporous ternary metal oxide $Cu_xCo_{3-x}O_4$ (x=0.5, 1) microcubes exhibit much better performance for CO oxidation than pure Co_3O_4 , which can be mainly attributed to their larger specific surface areas, stronger reducibility, and the synergistic effect of two active metal oxide components.

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Introduction

CO oxidation over solid catalysts has been extensively studied, not only with regard to its value as a classical probe reaction for fundamental studies but also due to its great importance in practical applications, such as air cleanup, gas sensors for detection, automotive exhaust gas treatment and CO elimination from the hydrogen feed for proton-exchange membrane fuel cells.1-5 It has been reported that noble metals such as Pt, Au, and Rh are excellent CO oxidation catalysts. 2,6-11 Platinum group metals (PGMs) are some of the earliest catalysts and are still widely employed in automotive emission control and cleaning of gas originating from the petrochemical industry since Langmuir's pioneering work.^{6,7,12} Haruta et al. discovered that oxide-supported Au catalysts prepared through a wet chemistry method exhibit an outstanding high activity for CO oxidation even at cryogenic temperatures. 13 Recently, a subnano Rh/TiO2 catalyst with an unexpectedly high performance of total CO conversion at -50 °C was discovered by Guan et al. 10 The excellent properties of this catalyst can be ascribed to the facile formation of Rh-O-O-Ti superoxide easily reacting with the adsorbed CO species on TiO₂ sites.

Although these noble metals exhibit high activities for CO oxidation, the high cost and limited supply constrain their commercial applications.14 Thus attention has been focused on searching for "noble metal-free" catalysts with excellent catalytic properties. 15-21 Co₃O₄, one kind of the less-expensive alternatives to the noble metal-based catalysts, has been reported to be one of the most efficient catalysts for carbon monoxide oxidation due to its excellent CO adsorption strength, low barrier of CO reacting with lattice O, and remarkable redox properties. 14,15,18,22 A pioneering work on Co₃O₄ for CO oxidation reported by Xie et al. revealed that the Co₃O₄ nanorods demonstrate a catalytic activity towards CO oxidation at a temperature of -77 °C.23 The superior performance of the catalyst can be attributed to the predominantly exposed (110) planes with richness of active Co3+ sites. A recent theoretical work found that Co³⁺ has a higher activity than Co²⁺ in the Co₃O₄ for CO oxidation because: (i) Co³⁺ binds CO molecule strongly compared to Co2+, (ii) a high barrier exists between lattice O linked to Co²⁺ and CO adsorbed on Co²⁺.²² Therefore, many works are focusing on preparing special nanostructural Co₃O₄ predominately exposing active faces with more Co³⁺ cations to enhance their catalytic activities for CO oxidation. However, these active faces still contain many Co²⁺ cations which have lower activities for CO oxidation.24-26 So, replacing the inactive Co²⁺ with other active divalent cations to prepare Cobased ternary metal oxides provides an efficient way to further enhance the catalytic performance of Co₃O₄. On this point, Cu²⁺ can be a promising candidate for replacing Co2+ in the Co3O4 not only due to its high activities for CO oxidation, but also owning to the similar ionic radius with Co2+ which allows the easy formation of Co/Cu ternary metal oxides.27-29

MOFs are a class of porous functional materials constructed by metal ions and bridging ligands.³⁰⁻³³ Owing to its adjustable

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cavities and flexible structures, various metal oxides with specific morphologies and interconnected pores have been fabricated by thermal decomposition of MOFs under suitable calcination conditions, expecting to improve their performance in specific applications.³⁴⁻⁴² Among them, ternary metal oxides have gained increasing considerations due to the structure merits and synergetic effect of multiple components. Therefore, bimetallic organic frameworks are considered highly desirable to facile synthesis of the ternary metal oxides. However such effort is still scarce because of the challenge in morphology and composition control during the incorporation of different secondary metal nodes into MOFs.39 Divalent metal formate frameworks (MFFs) of $[CH_3NH_3][M(HCOO)_3]$ (M = Mn, Fe, Ni, Co, Zn) could be promising MOFs to tackle this challenge due to the same six-connected $(4^{12} \cdot 6^3)$ nodes of the octahedral (MO₆) metal ions within the framework.43 Herein, we propose a facial one-step synthesis of bimetallic formate frameworks [CH₂NH₃] $[M(HCOO)_3]$ (M = Co/Cu) (Co/Cu-MFFs) and its derivation of ternary oxides $Cu_xCo_{3-x}O_4$ (x = 0, 0.5, 1, 1.5) microcubes. The Cu_xCo_{3-x}O₄ microcubes are constructed by numerous interconnected nanoparticles with uniform distribution of cobalt and copper species. In addition, the Cu/Co atomic ratio can be controlled exactly through adjusting the compositions of the growth precursor solutions. Owing to its larger specific surface area, stronger reducibility and the synergistic effect of two metal oxide components, the mesoporous Cu_rCo_{3-r}O₄ with moderate Cu/Co atomic ratio exhibit better performance for CO oxidation reaction than pure Co₃O₄.

Experimental 2.

Synthesis of Co/Cu-MFFs precursors

25 mL of ethanol was mixed with 12 mmol HCOOH, 8 mmol CH₃NH₂ (30-33% in methanol) and 0.5 g polyvinylpyrrolidone (PVP K-30) to get a colorless solution. Then a transparent solution containing 1 mmol mixed salts with different Cu/Co atomic ratios (0, 1:5, 1:2, 1:1) and 0.5 g PVP K-30 were dropped into the above solution in 10 minutes to obtain a pink colloidal suspension. The whole reaction process was maintained at room temperature with magnetic stirring. One hour later, the colloidal suspension was aged for one day at ambient temperature without any interruption. The resulting pink precipitates were centrifuged and washed several times with ethanol, finally dried at 60 °C for 6 hours in a vacuum oven and noted as [CH₃NH₃] $[Co(HCOO)_3]$, $[CH_3NH_3][Cu_{1/6}Co_{5/6}(HCOO)_3]$, $[CH_3NH_3][Cu_{1/6}Co_{5/6}(HCOO)_3]$ $_3\text{Co}_{2/3}(\text{HCOO})_3$, [CH $_3\text{NH}_3$][Cu $_{1/2}\text{Co}_{1/2}(\text{HCOO})_3$] respectively.

2.2. Synthesis of $Cu_xCo_{3-x}O_4$ (x = 0, 0.5, 1, 1.5) microcubes

The porous ternary metal oxides were fabricated in muffle burner by annealing these prepared Co/Cu-MFFs precursors at 350 °C for 2 hours, at a heating rate of 1 °C min⁻¹ and noted as Co₃O₄, Cu_{0.5}Co_{2.5}O₄, CuCo₂O₄, Cu_{1.5}Co_{1.5}O₄, respectively.

2.3. Characterizations

Powder XRD patterns were recorded on a Bruker D8 advance diffractometer with Cu K_{α} radiation over the 2θ range of 10–80°.

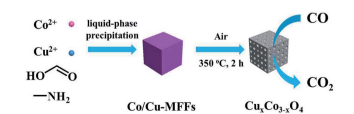
Thermogravimetric analyses (TGA) were carried out in a Netzsch SDT449F3 thermal analyzer in air atmosphere with a heating rate of 5 °C min⁻¹. The SEM and TEM images of the prepared samples were obtained through field emission scanning electron microscopy (FESEM, Hitachi S-4800 microscope) and transmission electron microscopy (TEM, JEOL, JEM-2100F). The chemical composition and elemental distribution of the prepared catalysts were examined by Energy-dispersive X-ray spectroscope (EDX) attached to the FESEM instrument. ICP-OES was used to determine the Cu/Co atomic ratio of Cu_xCo_{3-x}O₄. The X-ray photoelectron spectroscopy (XPS) spectra were collected on Thermo ESCALAB 250 Xi spectrometer. N₂ adsorption-desorption isotherm were obtained by a Micromeritics ASAP 2020 analyzer at 77 K. H₂temperature-programmed reduction (H2-TPR) experiments were performed under a 10 vol% H₂/Ar mixture with a flow rate of 50 mL min⁻¹ over 60 mg of catalyst by a Micromeritics Chemisorb 2920 apparatus and the temperature was increased from ambient temperature to 800 °C at a ramp rate of 1 °C min⁻¹. Prior to each analysis, the catalysts were purged in a flow of pure argon at 200 °C for 2 hours to remove traces water.

2.4. Catalytic performance measurements

The catalytic tests of the Cu_xCo_{3-x}O₄ microcubes for CO oxidation reaction were performed in a fixed quartz tubular reactor. The reactor was charged with 50 mg of the catalysts without pretreatment. The reaction temperature was detected by a thermocouple under catalyst bed. The mixed feed gas consisted of 1 vol% CO, 20 vol% O2, 79 vol% N2 was passed through the reactor at a flow rate of 50 mL min⁻¹, corresponding to a gas hourly space velocity (GHSV) of 60 000 mL g⁻¹ h⁻¹. The components of the gas were examined by a gas chromatograph (Agilent GC7890). Two catalytic test data points at each temperature were collected and the average values of them were reported.

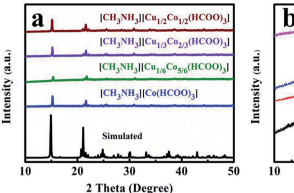
3. Results and discussion

The synthesis process of mesoporous Cu_xCo_{3-x}O₄ microcubes is shown in Scheme 1. Firstly, the Co/Cu-MFFs precursors with different Cu/Co atomic ratios were synthesized through a onestep liquid-phase precipitation method under room temperature. Then, the Cu_xCo_{3-x}O₄ microcubes with porous structure were obtained through thermal decomposition of the precursors under air. The phase and purity of the Co/Cu-MFFs precursors and Cu_xCo_{3-x}O₄ were examined by PXRD (Fig. 1a). All strong peaks in the XRD patterns of each Co/Cu-MFFs



Scheme 1 Schematic illustration of the synthesis of Co/Cu-MFFs precursors and its derived mesoporous Cu_xCo_{3-x}O₄ microcubes for CO oxidation reaction.

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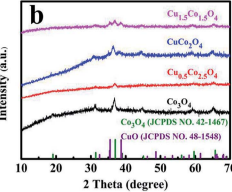


Fig. 1 XRD patterns of Co/Cu-MFFs precursors (a) and Cu_xCo_{3-x}O₄ microcubes (b).

precursor are well matched with the simulated single crystal structure of [CH₃NH₃][Co(HCOO)₃] according to the reported crystal structure data,44,45 suggesting the successful incorporation of Cu²⁺ into Co/Cu-MFFs. Fig. S1† shows the TGA curves of as-prepared precursors, indicating that the weight residual of all samples becomes constant when the temperature rises to 300 °C. During the calcination, all samples exhibit similar twostep weight loss. The first weight loss could be relate to the removal of amine and one formate molecule per formula unit. The second weight loss could be due to the decomposition of residual organic components. Fig. 1b shows that the corresponding diffraction profiles of prepared ternary oxides Cu_x-Co_{3-x}O₄ are in agreement with the standard CuCo₂O₄ (JCPDS no. 01-1155) or pure Co₃O₄ (JCPDS no. 42-1467). CuCo₂O₄ can be treated as Co₃O₄ with Co²⁺ replaced by Cu²⁺ and therefore these two oxides have almost same XRD patterns. Meanwhile, two weak peaks corresponding to CuO (002), (111) (JCPDS no. 48-1548) can be seen in the XRD patterns of CuCo2O4 and Cu_{1.5}Co_{1.5}O₄, suggesting that a few Cu²⁺ cations did not enter into the lattice of Co₃O₄ successfully. These two peaks become obvious with the increase of Cu-doping amount, indicating that the excess amounts of Cu²⁺ cations in the Co/Cu-MFFs precursors lead to a significant phase segregation of the oxides.

The SEM images in Fig. S2a and S2b† reveal that the asprepared [CH₃NH₃][Co(HCOO)₃] and [CH₃NH₃][Cu_{1/3}Co_{2/} 3(HCOO)3] have similar cubic shapes with smooth surfaces and some macropores about 0.6 µm can be observed clearly on the partial surfaces of these two precursors. Meanwhile, the size of [CH₃NH₃][Cu_{1/3}Co_{2/3}(HCOO)₃] microcubes is about 1-2 μm, which is smaller than that of [CH₃NH₃][Co(HCOO)₃], mainly about 2-3 µm. We can infer accordingly that the introduced Cu²⁺ does not have great influences on the morphology of precursors. In order to study the effect of PVP K-30 in the preparation of Co/Cu-MFFs precursors, the morphology of [CH₃NH₃][Co(HCOO)₃] prepared in the absence of PVP K-30 was also characterized. As shown in Fig. S3,† the [CH3NH3] [Co(HCOO)₃] without adding PVP K-30 still retains the cubic morphology but many microbes agglomerate together. So we can infer that PVP K-30 functions here mainly as stabilizing agent to prevent agglomeration of the microcubes. The similar functions have been reported by many other reports. 46,47 The

SEM images of Cu_xCo_{3-x}O₄ easily prepared by annealing precursors are shown in Fig. 2a-d. The cubic morphology of Cu_xCo_{3-x}O₄ reveals that all the Cu_xCo_{3-x}O₄ almost preserved the original cubic morphology of as-prepared precursors. Meanwhile, a great number of small nanoparticles on the rough surface of cubic Cu_xCo_{3-x}O₄ can be seen clearly in the SEM images, which confirms the successful formation of porous structure. The chemical composition and elemental distribution of the microcubes were further characterized by SEM-EDX. The elemental mapping images (Fig. 2e and S4†) further confirm that all the ternary metal oxides are with similar microcubic morphologies and the uniform distribution of O, Cu and Co. The metal element ratios of Cu and Co in the oxides are similar with those ratios in the growth precursor solution as shown in Table S1 and Fig. S5,† which shows that the compositions of Cu_xCo_{3-x}O₄ can be controlled exactly by adjusting the synthesis of $[CH_3NH_3][Cu_{1/3}Co_{2/3}(HCOO)_3]$. The high-resolution TEM image of CuCo₂O₄ microcube is shown in Fig. 2f, the interplanar distances of 0.23 nm can be indexed to the (111) plane of CuO, whereas the interplanar distance of 0.24 nm is correspondence to the (311) plane of CuCo₂O₄. The interlaced boundaries marked with white circle demonstrate the high interdispersion of the CuO and CuCo₂O₄.

Some related mechanisms at the molecular level of Co₃O₄ for CO oxidation suggest that gas phase CO chemisorbs preferably on the exposed Co3+, then reacts with an oxygen atom linked to the active Co3+ site resulting in CO2 and an oxygen vacancy formation.24,48 Therefore, the amount of the exposed Co3+ cations on the Co₃O₄ surfaces is responsible for catalytic activity.49 Herein, the XPS analysis was further carried out to examine the surface chemical compositions and elemental states of Cu_rCo_{3-r}O₄ microcubes. The Co2p XPS spectra profiles (Fig. 3a) constructed with two main peaks centering at about 779.5 and 794.6 eV, corresponding to the $Co2p_{3/2}$ and $Co2p_{1/2}$ respectively.37,50,51 The relative percentage of Co3+ and Co2+ was calculated through the fitted curves of CuxCo3-xO4, and was plotted as functions of the Cu/Co atomic ratio in the solution of preparing precursors (Fig. 3b). We can find that the ratio of Co^{2+/}Co³⁺ on the surface of Cu_rCo_{3-r}O₄ first drastically decreases and then almost remains unchanged with the increasing Cu/Co atomic ratio. That decline of Co²⁺/Co³⁺ atomic

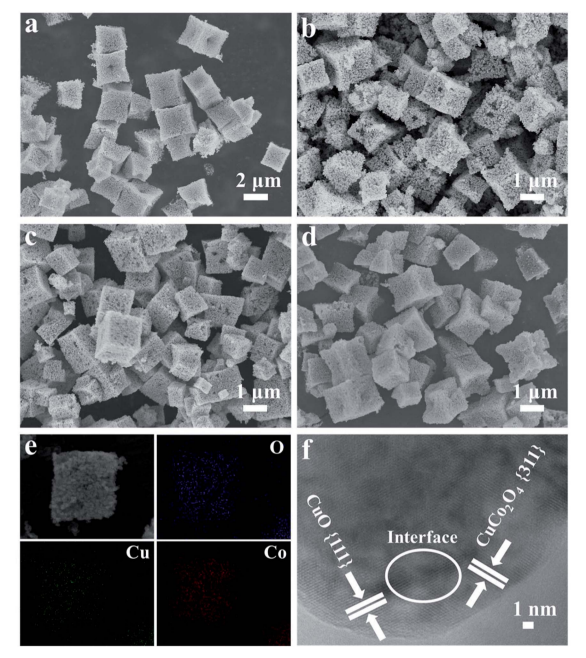


Fig. 2 SEM images of Co₃O₄ (a), Cu_{0.5}Co_{2.5}O₄ (b), CuCo₂O₄ (c), Cu_{1.5}Co_{1.5}O₄ (d), SEM-EDX mapping images of CuCo₂O₄ (e), TEM images of $CuCo_2O_4$ (f).

ratio should be originated from the Cu²⁺ substitution for Co²⁺ in $Cu_xCo_{3-x}O_4$. The further steadiness might be ascribed to many Cu²⁺ leading to the formation of CuO rather than substitution for Co²⁺, which can also be inferred from the XRD results.

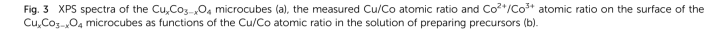
The porous structures of Co/Cu-MFFs derived CuCo₂O₄ and pure Co₃O₄ were further examined by measuring sorption isotherms of nitrogen at 77 K. It can be observed in Fig. 4a that both of the oxides present a type-IV adsorption isotherm with a significant hysteresis loop, denoting that they are mesoporous solids. The specific surface areas of Co₃O₄ and CuCo₂O₄ microcubes were calculated to be 25.59 and 29.47 m² g⁻¹ through Brunauer-Emmett-Teller (BET) method. In addition, the corresponding Barrett-Joyner-Halenda (BJH) pore size distribution plots (Fig. S6†) show the main pore size distribution of Co₃O₄ and CuCo₂O₄ are 0.71-1.58 nm and 16.15-20.73 nm, respectively.

To directly evaluate the catalytic performance of the Co/Cu-MFFs derived ternary oxides for CO oxidation reaction, 50 mg of the mesoporous Cu_xCo_{3-x}O₄ microcubes without any pretreatments were put into a fixed quartz glass reactor respectively. As shown in Fig. 4b, with the increasing amount of Cu/Co atomic ratio in the catalysts, the catalytic activities of Cux-Co_{3-x}O₄ for CO oxidation reaction firstly increase and then decrease. Among them, CuCo2O4 exhibits the highest activity

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Co3+ Co2+ 0.65 Cu/Co molar ratio from EDX Co₃O₄ Co₃O₄ 1.0 0.60 Intensity (a.u.) Cu_{0.5}Co_{2.5}O₄-1:5 Cu_{0.5}Co_{2.5}O₄ 0.8 0.55 0.6 CuCo,O4 0.50 0.4 CuCo2O4 Cu_{1.5}Co_{1.5}O 0.2 Cu_{1.5}Co_{1.5}O

800



0.0

0.0

0.2

0.4

Cu/Co atomic ratio in the precursor

0.6

0.8

1.0

with 100% CO conversion rate at 120 °C, superior or comparable to other transition oxides as CO oxidation catalysts (Table S2†). These results indicate that using more active Cu²⁺ to replace the inactive Co²⁺ in the lattice of Co₃O₄ could be a novel way to enhance the catalytic performance for CO oxidation. Meanwhile, the Cu_{1.5}Co_{1.5}O₄ has the lowest activity and the temperature of CO complete conversion is as high as 170 °C, which might be due to the richness of less-active CuO phase. The CO oxidation conversions of without materials and with 50 mg of precursors performed under different temperatures were shown

780

790

Binding energy (eV)

in Fig. S7,† the results indicated that there was no catalytic activity in the absence of catalyst and the Co/Cu-MFFs exhibited extremely low catalytic activity of CO oxidation. We also investigated the stabilities of pure Co₃O₄ and CuCo₂O₄ under the similar conditions at 140 and 130 °C, respectively (Fig. 4c). Both samples exhibit good stable performance that retains CO complete conversion within 30 hours. Based on these results, we can infer that the enhanced catalytic activity of CurCo_{3-r}O₄ can be attributed to the structure and component merits achieved through the incorporation of new secondary copper nodes

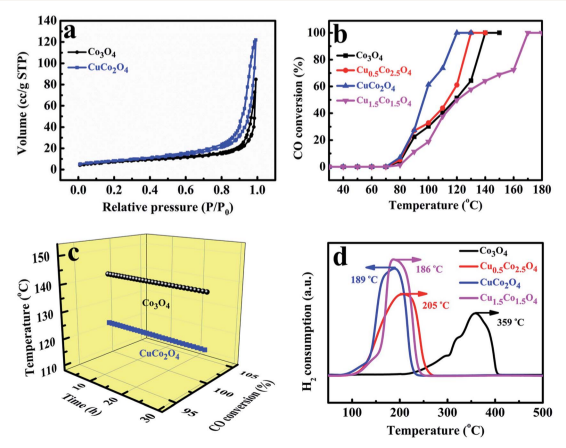


Fig. 4 Nitrogen adsorption – desorption isotherms of Co_3O_4 and $CuCo_2O_4$ (a), the catalytic activities of $Cu_xCo_{3-x}O_4$ for CO oxidation (b), stability of Co₃O₄ at 140 °C and CuCo₂O₄ at 120 °C (c), H₂-TPR profiles of Cu_xCo_{3-x}O₄ (d).

in the Co/Cu-MFFs precursors. In addition, pure ${\rm Co_3O_4}$ presents a good activity for CO oxidation with a CO complete conversion temperature of 140 °C, which is 30 °C lower than that reported by Zhang $et~al.^{52}$ The higher catalytic activity might be ascribed to the larger specific surface area and the lower inactive ${\rm Co^{2^+}}$ concentration in the surface of ${\rm Co_3O_4}$ according to the BET and

To further investigate the synergetic effects of Cu and Co species, the temperature-programmed reductions were performed using a stream of diluted H₂ (H₂/Ar₂, 1:9 v/v). As shown in Fig. 4d, only a single peak at 359 °C is observed in pure Co₃O₄ microcubes reduction profile, which can be explained by the direct reduction of cobalt ions into metallic Co in one step for the large particles of Co₃O₄.53 With the amount of Cu²⁺ increasing in catalysts, the prominent peaks of Cu_{0.5}Co_{2.5}O₄, CuCo₂O₄ and Cu_{1.5}Co_{1.5}O₄ shift towards lower temperature to 205, 189 and 186 °C, respectively, which means that the reducibility of the catalysts was markedly promoted when the Cu/Co atomic ratio increased. Such enhanced reduction may be attributed to the strong Co-Cu interaction originated from the intimate contact and the good interdispersion of the CuO and Co₃O₄, which possibly leads to a junction of their band levels in the solid solution or the mixed oxidations.⁵⁴ Such strong interactions also have been found for copper-zinc chromite catalysts and CuO-ZnO system.55,56

4. Conclusion

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XPS results (Table S3†).52

In this work, we have proposed a novel and simple method to prepare Co-based ternary metal oxides Cu_xCo_{3-x}O₄ with cube-like morphology and studied the effect of replacing Co²⁺ with Cu²⁺ on the catalytic activity of Co₃O₄ towards CO oxidation. The Cu_x-Co_{3-x}O₄ microcubes with controlled composition were prepared by one step pyrolysis of Co/Cu-MFFs, in which the Cu/Co atomic ratio can be tuned through adjusting the compositions of growth precursor solution. The obtained mesoporous Cu_xCo_{3-x}O₄ have enhanced catalytic properties for CO oxidation compared to the pure porous Co₃O₄ when the atomic ratio of Cu/Co is no more than 1/2 in the solution of preparing precursors. The larger specific surface areas and stronger reducibility resulted from the introduced new secondary Cu metal nodes into Co/Cu-MFFs which may play important roles in improving catalytic performance. We believe that the synthetic strategy can also be extended to obtain other ternary metal oxides with different structures by controlling the morphology and the compositions of the bimetal-formate frameworks, which might have a higher catalytic performance or exhibit excellent properties in other applications.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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References

- 1 H.-J. Freund, G. Meijer, M. Scheffler, R. Schlögl and M. Wolf, *Angew. Chem., Int. Ed.*, 2011, **50**, 10064–10094.
- M. Haruta, S. Tsubota, T. Kobayashi, H. Kageyama,
 M. J. Genet and B. Delmon, *J. Catal.*, 1993, 144, 175–192.
- 3 J.-C. Ding, H.-Y. Li, T.-C. Cao, Z.-X. Cai, X.-X. Wang and X. Guo, *Solid State Ionics*, 2017, **303**, 97–102.
- 4 K. M. Adams and G. W. Graham, Appl. Catal., B, 2008, 80, 343–352.
- 5 P. Landon, J. Ferguson, B. E. Solsona, T. Garcia, A. F. Carley, A. A. Herzing, C. J. Kiely, S. E. Golunski and G. J. Hutchings, *Chem. Commun.*, 2005, 3385–3387.
- 6 J. Lin, X. Wang and T. Zhang, *Chin. J. Catal.*, 2016, 37, 1805–1813.
- 7 B. Liu, Y. Liu, H. Hou, Y. Liu, Q. Wang and J. Zhang, Catal. Sci. Technol., 2015, 5, 5139–5152.
- 8 L. W. Guo, P. P. Du, X. P. Fu, C. Ma, J. Zeng, R. Si, Y. Y. Huang, C. J. Jia, Y. W. Zhang and C. H. Yan, *Nat. Commun.*, 2016, 7, 13481.
- 9 B. Qiao, J. Lin, A. Wang, Y. Chen, T. Zhang and J. Liu, *Chin. J. Catal.*, 2015, **36**, 1505–1511.
- 10 H. Guan, J. Lin, B. Qiao, X. Yang, L. Li, S. Miao, J. Liu, A. Wang, X. Wang and T. Zhang, Angew. Chem., Int. Ed., 2016, 55, 2820–2824.
- 11 H. Guan, J. Lin, L. Li, X. Wang and T. Zhang, Appl. Catal., B, 2016, 184, 299–308.
- 12 I. Langmuir, Trans. Faraday Soc., 1992, 17, 621-654.
- 13 M. Haruta, N. Yamada, T. Kobayashi and S. Iijima, *J. Catal.*, 1989, **115**, 301–309.
- 14 L. F. Liotta, H. Wu, G. Pantaleo and A. M. Venezia, *Catal. Sci. Technol.*, 2013, **3**, 3085–3102.
- 15 X. Wang, W. Zhong and Y. Li, Catal. Sci. Technol., 2015, 5, 1014–1020.
- 16 Z. Zhao, X. Lin, R. Jin, G. Wang and T. Muhammad, *Appl. Catal.*, *B*, 2015, **115–116**, 53–62.
- 17 Y. Su, Z. Tang, W. Han, Y. Song and G. Lu, *Catal. Surv. Asia*, 2015, **19**, 68–77.
- 18 C. J. Jia, M. Schwickardi, C. Weidenthaler, W. Schmidt, S. Korhonen, B. M. Weckhuysen and F. Schuth, *J. Am. Chem. Soc.*, 2011, 133, 11279–11288.
- 19 A. Biabani-Ravandi and M. Rezaei, *Chem. Eng. J.*, 2012, **184**, 141–146
- 20 T. Cwele, N. Mahadevaiah, S. Singh and H. B. Friedrich, *Appl. Catal.*, *B*, 2016, **182**, 1–14.
- 21 Y. Yu, T. Takei, H. Ohashi, H. He, X. Zhang and M. Haruta, *J. Catal.*, 2009, **267**, 121–128.
- 22 H.-F. Wang, R. Kavanagh, Y.-L. Guo, Y. Guo, G. Lu and P. Hu, J. Catal., 2012, 296, 110–119.
- 23 X. Xie, Y. Li, Z. Q. Liu, M. Haruta and W. Shen, *Nature*, 2009, 458, 746–749.
- 24 J. Jansson, A. E. C. Palmqvist, E. Fridell, M. Skoglundh, L. Österlund, P. Thormählen and V. Langer, *J. Catal.*, 2002, 211, 387–397.
- 25 P. Broqvist, I. Panas and H. Persson, *J. Catal.*, 2002, **210**, 198–206.

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26 F. Grillo, M. M. Natile and A. Glisenti, *Appl. Catal., B*, 2004, **48**, 267–274.

- 27 M. Y. Guo, F. Liu, J. Tsui, A. A. Voskanyan, A. M. C. Ng, A. B. Djurišić, W. K. Chan, K.-Y. Chan, C. Liao, K. Shih and C. Surya, *J. Mater. Chem. A*, 2015, 3, 3627–3632.
- 28 R. Zhang, L. Hu, S. Bao, R. Li, L. Gao, R. Li and Q. Chen, *J. Mater. Chem. A*, 2016, **4**, 8412–8420.
- 29 M. Zhou, L. Cai, M. Bajdich, M. García-Melchor, H. Li, J. He, J. Wilcox, W. Wu, A. Vojvodic and X. Zheng, ACS Catal., 2015, 5, 4485–4491.
- 30 E. A. Dolgopolova, A. J. Brandt, O. A. Ejegbavwo, A. S. Duke, T. D. Maddumapatabandi, R. P. Galhenage, B. W. Larson, O. G. Reid, S. C. Ammal, A. Heyden, M. Chandrashekhar, V. Stavila, D. A. Chen and N. B. Shustova, *J. Am. Chem. Soc.*, 2017, 139, 5201–5209.
- 31 L. Chen, R. Luque and Y. Li, *Chem. Soc. Rev.*, 2017, **46**, 4614–4630.
- 32 X. Yang and Q. Xu, Cryst. Growth Des., 2017, 17, 1450-1455.
- 33 L. Zhu, X. Q. Liu, H. L. Jiang and L. B. Sun, Chem. Rev., 2017, 117, 8129–8176.
- 34 F. Wang, X. Wang, D. Liu, J. Zhen, J. Li, Y. Wang and H. Zhang, *ACS Appl. Mater. Interfaces*, 2017, **6**, 22216–22223.
- 35 Y. Zhu, C. Cao, J. Zhang and X. Xu, J. Mater. Chem. A, 2015, 3, 9556–9564.
- 36 L. Shen, L. Yu, X.-Y. Yu, X. Zhang and X. W. D. Lou, Angew. Chem., Int. Ed., 2015, 54, 1868–1872.
- 37 J. Wei, Y. Feng, Y. Liu and Y. Ding, *J. Mater. Chem. A*, 2015, 3, 22300–22310.
- 38 L. Han, X.-Y. Yu and X. W. D. Lou, *Adv. Mater.*, 2016, 28, 4601–4605.
- 39 H. Li, M. Liang, W. Sun and Y. Wang, Adv. Funct. Mater., 2016, 26, 1098–1103.
- 40 R. R. Salunkhe, Y. V. Kaneti and Y. Yamauchi, *ACS Nano*, 2017, **11**, 5293–5308.
- 41 Z. Yu, Y. Bai, Y. Liu, S. Zhang, D. Chen, N. Zhang and K. Sun, *ACS Appl. Mater. Interfaces*, 2017, 9, 31777–31785.

- 42 S. Zheng, X. Li, B. Yan, Q. Hu, Y. Xu, X. Xiao, H. Xue and H. Pang, *Adv. Energy Mater.*, 2017, 7, 1602733.
- 43 Z. Wang, K. Hu, S. Gao and H. Kobayashi, *Adv. Mater.*, 2010, **22**, 1526–1533.
- 44 L. C. Gomez-Aguirre, B. Pato-Doldan, J. Mira, S. Castro-Garcia, M. A. Senaris-Rodriguez, M. Sanchez-Andujar, J. Singleton and V. S. Zapf, *J. Am. Chem. Soc.*, 2016, 138, 1122–1125.
- 45 B. Pato-Doldan, L. C. Gomez-Aguirre, A. P. Hansen, J. Mira, S. Castro-Garcia, M. Sanchez-Andujar, M. A. Senaris-Rodriguez, V. S. Zapf and J. Singleton, *J. Mater. Chem. C*, 2016, 4, 11164–11172.
- 46 X. Huang and N. Zheng, *J. Am. Chem. Soc.*, 2009, **131**, 4602–4603.
- 47 J. Yin, J. Wang, M. Li, C. Jin and T. Zhang, *Chem. Mater.*, 2012, 24, 2645–2654.
- 48 Y. Xie, F. Dong, S. Heinbuch, J. J. Rocca and E. R. Bernstein, *Phys. Chem. Chem. Phys.*, 2010, 12, 947–959.
- 49 A. Alvarez, S. Ivanova, M. A. Centeno and J. A. Odriozola, Appl. Catal., A, 2012, 431–432, 9–17.
- 50 G. Li, L. Li, Y. Li and J. Shi, New J. Chem., 2015, 39, 1742-1748.
- 51 J. Zhu and Q. Gao, Microporous Mesoporous Mater., 2009, 124, 144–152.
- 52 C. Zhang, L. Zhang, G.-C. Xu, X. Ma, Y.-H. Li, C.-Y. Zhang and D.-Z. Jia, *New J. Chem.*, 2017, 41, 1631–1636.
- 53 J.-Y. Luo, M. Meng, X. Li, X.-G. Li, Y.-Q. Zha, T.-D. Hu, Y.-N. Xie and J. Zhang, *J. Catal.*, 2008, **254**, 310–324.
- 54 G. Fierro, M. Lo Jacono, M. Inversi, R. Dragone and P. Porta, *Top. Catal.*, 2000, **10**, 39–48.
- 55 G. L. Castiglioni, A. Vaccari, G. Fierro, M. Inversi, M. Lo Jacono, G. Minelli, I. Pettiti, P. Porta and M. Gazzano, *Appl. Catal.*, *A*, 1995, 123, 123–144.
- 56 G. Fierro, M. Lo Jacono, M. Inversi, P. Porta, F. Cioci and R. Lavecchia, *Appl. Catal.*, *A*, 1996, 137, 327–348.